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- 1. A method for labeling genetic material, the method comprising:
- disrupting cells so as to liberate genetic material contained in the cells;
- b) contacting the genetic material to a column in a manner to cause the genetic material to become immobilized to the column;
 - labeling the immobilized genetic material within the column; and c)
 - d) eluting the labeled material from the column.
 - 2. A method for manipulating genetic material, the method comprising:
- disrupting cells so as to liberate genetic material contained in the cells:
- contacting the genetic material to a column in a manner to cause the genetic material to become immobilized to the column;
 - labeling the immobilized genetic material; and C)
- dY eluting the labeled material from the column wherein the step of labeling the genetic material further comprises maintaining the column at a temperature of between 45 °C and 100 °C.
 - 5. A method for manipulating genetic material, the method comprising:
- disrupting cells so as to liberate genetic material contained in the cells;
- b) contacting the genetic material to a column in a manner to cause the genetic material to become immobilized to the column;
 - labeling the immobilized genetic material; and c)
 - d) eluting the labeled praterial from the column wherein the step of labeling the genetic material comprises:
- contacting double-stranded nucleic acid molecules of the genetic material with radical-generating complexes for a time and at concentrations sufficient to produce free-aldehyde moieties;
- reacting the aldehyde moieties with amine to produce a condensation f) product; and,
 - contacting the condensation product with a chromophore.
 - 8. A two-buffer process for labeling genetic material, the process comprising:
 - contacting cells containing the genetic material to a silica column;
- b) creating a first fraction of cell detritus and a second fraction containing the genetic material;
 - confining the genetic material to the column; c)
 - removing the cell detritus; d)
 - subjecting the genetic material to radicals so as to produce reactive e)



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aldehyde groups on the genetic material; and

- f) attaching chromophore to the genetic material while the material resides in the column.
- 9. A two-buffer process for manipulating genetic material, the process comprising:
 - a) contacting cells containing the genetic material to a silica column;
- b) creating a first fraction of cell detritus and a second fraction containing the genetic material;
 - c) confining the genetic material to the column;
 - d) removing the cell detritus;
- e) subjecting the genetic material to radicals so as to produce reactive aldehyde groups on the genetic material; and
 - f) attaching chromophore to the genetic material wherein the genetic material is contacted with radical in aerobic conditions.
- 10. A two-buffer process for manipulating genetic material, the process comprising:
 - a) contacting cells containing the genetic material to a silica column;
- b) creating a first fraction of cell detritus and a second fraction containing the genetic material;
 - c) confining the genetic material to the column;
 - d) removing the cell detritus;
- e) subjecting the genetic material to radicals so as to produce reactive aldehyde groups on the genetic material; and
 - f) attaching chromophore to the genetic material wherein the genetic material is contacted with radical in anaerobic conditions.
- 13. A two-buffer process for manipulating genetic material, the process comprising:
 - a) contacting cells containing the genetic material to a silica column;
- b) creating a first fraction of cell detritus and a second fraction containing the genetic material;
 - c) confining the genetic material to the column;
 - d) removing the cell detritus;
- e) subjecting the genetic material to radicals so as to produce reactive aldehyde groups on the genetic material; and
- f) attaching chromophore to the genetic material wherein the two buffers comprise a first buffer to lyse the cells and a second buffer to attach the genetic material to the column.



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- 19. The process as recited in claim 8 wherein the temperature is maintained at between 30 °C and 100 °C.
- 20. The method as recited in claim 2 wherein the column comprises a means for subjecting the silica to pressure.
- 21. The method as recited in claim 1 wherein the step of labeling the genetic material comprises:
- a) contacting nucleic acid molecules of the genetic material with radicalgenerating complexes for a time and at concentrations sufficient to produce freealdehyde moieties;
- b) reacting the aldehyde moieties with amine to produce a condensation product; and
 - c) contacting the condensation product with a chromophore.
- 22. The method as recited in claim 21 wherein the step of contacting the condensation product with a chromophore further comprises reducing the condensation product and cross-linking the reduced condensation product with the chromophore in one reaction step.
- 23. The process as recited in claim 9 wherein the genetic material is bound to chromophore in aerobic conditions.
- 24. The process as recited in claim 10 wherein the genetic material is bound to chromophore in anaerobic conditions.
- 25. The process as recited in claim 8 wherein the two buffers comprise a first buffer to lyse the cells and a second buffer to attach the genetic material to the column.

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